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(54) Title: IN VITRO ANTIBODY AFFINITY MATURATION USING ALANINE SCANNING MUTAGENESIS

(57) Abstract

A method of mutagenizing antibodies to produce modified antibodies, modified antibodies, DNA encoding the modified antibodies as well as diagnostic kits and pharmaceutical compositions comprising the antibodies or DNA are provided.

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TITLE OF THE INVENTION

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IN VITRO ANTIBODY AFFINITY MATURATION USING ALANINE SCANNING MUTAGENESIS

CROSS-RELATED TO OTHER APPLICATIONS

This is a continuation of U.S. Serial No. 08/206,076 filed March 4, 1994, now pending.

BRIEF DESCRIPTION OF INVENTION

A method of mutagenizing antibodies to produce 10 modified antibodies, modified antibodies, DNA encoding the modified antibodies as well as diagnostic kits and pharmaceutical compositions comprising the antibodies or DNA are provided. The method of the invention is a systematic means to achieve in vitro antibody maturation and uses alanine scanning mutagenesis. The 15 invention is particularly exemplified with a set of single chain Fv (scFv) antibodies obtained by this technique. The resulting antibodies are directed against the V3 loop of HIV gp120, and show altered off-rates against the antigen compared to the starting antibody. Of particular interest are the altered antibodies which 20 show improved (slower) off-rates to the antigen. Observed improvements have been as high as eleven-fold over wild-type.

SUMMARY OF THE INVENTION

A method of mutagenizing antibodies to produce modified antibodies, modified antibodies, DNA encoding the modified antibodies as well as diagnostic kits and pharmaceutical compositions comprising the antibodies or DNA are provided.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1. Alanine-Scanning Mutagenesis. Each of the 27 amino acids in VH CDR3 of scFv P5Q was converted to alanine by site-directed mutagenesis. *E. coli* clones were induced to express scFv with IPTG. Single chain Fv, which is targeted to the periplasmic space by the fd phage gene3 signal sequence, was

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extracted with EDTA. Periplasmic extracts were analyzed by BIAcoreTM, which measures antibody-antigen affinity by surface plasmon resonance (Fägerstam, 1991), and off-rates determined against an HIV gp120 V3 loop peptide. Results of the alanine scan, relative to P5Q, fall into four classes: i) slower off-rate, ii) faster off-rate, iii) no binding, and iv) minor or no change in off-rate. Standard deviation is $\pm 25\%$.

Figure 2. Amino Acid Randomization: Position 107. Arginine at position 107 was mutated to all amino acids by site-directed mutagenesis. Single chain Fv extracts were analyzed by BIAcore. Percent change in off-rates is shown relative to P5Q.

Figure 3. Amino Acid Randomization: Position 111. Glutamic acid at position 111 was mutated to all amino acids by site-directed mutagenesis. Single chain Fv extracts were analyzed by BIAcore. Percent change in off-rates is shown relative to P5Q.

Figure 4. Amino Acid Randomization: Position 112. Aspartic acid at position 112 was mutated to all amino acids by site-directed mutagenesis. Single chain Fv extracts were analyzed by BIAcore. Percent change in off-rates is shown relative to P5Q.

Figure 5. Additive Effect of Combining Optimized Residues. A double mutant, containing the optimized residues, was constructed and analyzed by BIAcore. Percent change in off-rates is shown relative to P5Q.

Figure 6. Nucleotide and amino acid sequences of scFv P5O with c-myc tail.

DETAILED DESCRIPTION OF THE INVENTION

The gp120 V3 domain of human immunodeficiency virus-1 (HIV-1) is a disulfide-linked closed loop of approximately 30 amino acids. The loop, in either native or synthetic form, binds to and elicits anti-HIV-1 antibodies.

The present invention relates to modified antibodies and methods of making modified. The invention is exemplified with modified HIV-1 immunoglobulins and methods of making these

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modified HIV-1 immunoglobulins. The modified immunoglobulins of the present invention contain an altered complementary determining region 3 (CDR3) of HIV-1 neutralizing antibody.

The present invention also comprises a method of treating of preventing infection through the administration of a modified antibody to a suitable host. In one embodiment of the invention, the treatment or prevention of HIV infection through the administration of the modified HIV-1 immunoglobulin is described.

The present invention also comprises diagnostic kits useful for the detection or characterization of an antigen. Reagents for the kits may include DNA molecules encoding the modified antibodies or the modified antibodies or combinations thereof.

A method of mutagenizing antibodies to produce modified antibodies, modified antibodies, DNA encoding the modified antibodies as well as diagnostic kits and pharmaceutical compositions comprising the antibodies or DNA are provided. The method of the invention is a systematic means to achieve *in vitro* antibody maturation and uses alanine scanning mutagenesis. The invention is particularly exemplified with a set of single chain Fv (scFv) antibodies obtained by this technique. The resulting antibodies are directed against the V3 loop of HIV gp120, and show altered off-rates against the antigen compared to the starting antibody. Of particular interest are the altered antibodies which show improved (slower) off-rates to the antigen. Observed improvements have been as high as eleven-fold over wild-type.

Maturation was achieved through an alanine scan of complementary determining region 3 (CDR3) to identify positions critical to antigen binding. Critical positions were then randomized to identify amino acids that provided the slowest off-rates. Finally, clones were optimized through the combining of mutations.

The underlying principle of the method is the physical and chemical neutrality of alanine. Alanine is substituted throughout a stretch of amino acids, and its effects on binding (such as off-rate and on-rate) are evaluated using conventional methods. The number

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of positions likely to be identified in this manner is relatively small. Once identified, these key positions may be randomized to all amino acids to identify the best amino acid solution at the position. Because all manipulations and evaluations are conducted *in vitro*, physiological bias is limited.

Present methods of *in vitro* antibody maturation are essentially random procedures in which the researcher generates clones with amino acid substitutions and evaluates them. The problem is that the number of substitutions necessary for a thorough evaluation is extremely large. For example, if one were to evaluate all random substitutions in CDR3, a region typically twenty-five residues in length, one would have to examine 9-10²⁷ possibilities. This is beyond the capabilities of present technologies.

Alanine scanning maturation enables the rapid identification of residues most likely to be important in binding. Using the example of a twenty-five residue stretch cited above, only twenty-five substitutions would be necessary. From this initial screen, amino acid positions likely to be critical to binding may be identified. The critical residues may then be randomized to identify the amino acids that optimize binding. Using this method, scFv antibodies with dissociation rates greater than ten-fold slower than the original scFv have been created.

Previous work in *in vitro* antibody maturation used one of two general approaches. In one approach, PCR recombination is used to substitute all or part of the VH and VL genes into libraries of scFv clones. In the second approach, random mutations are made throughout a CDR region of a scFv clone by the use of degenerate oligonucleotides. In both cases, clones were expressed as a phage fd gene 3 fusion surface protein. Higher affinity clones were identified using a panning assay followed by clonal purification of the phage.

Each approach has drawbacks. The PCR method is cumbersome, limited to the sequences of the B cell population, is essentially random in nature, and may introduce unwanted mutations through the PCR recombination step. The randomization approach

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produces only a small fraction of the possible CDR changes. Neither approach allows immediate determination of changes in binding affinity because it is necessary to first generate an enriched population of suitable clones through panning. Both approaches detect only changes which result in improved binding; they do not identify positions for which the change weakened the binding. The latter class of change may include critical binding residues in which the appropriate amino acid solutions leads to improvement.

The method disclosed herein is systematic, thorough and unlikely to introduce unexpected or undesired mutations. All manipulations are done *in vitro*, which minimizes bias due to selection steps. Evaluation of clones is quantitative. In some cases, a key amino acid position may display poorer binding with alanine, but subsequent randomization may yield an amino acid solution which enables improved binding. Such mutations would not be detected by previous methods. Because the method of the present invention does not require phage expression for panning, the method can be used on scFVs, Fabs, and full length antibodies. Use is not restricted to a scFv for phage expression. Using the approach of the present invention, an anti-HIV V3 loop antibody was improved approximately eleven-fold.

Alanine scanning maturation of antibodies is a general method which may be used to improve binding of antibodies to their cognate antigens. The method has been used to identify critical residues in the scFv 447 which can be introduced into MAb447. Such changes may lead to significant improvement of the binding affinity of MAb447 against multiple species of HIV gp120 isolates. This improvement may increase the neutralization capability of the antibody, and significantly lower the effective dose.

Although the method and antibodies of the present invention are exemplified with scFv antibodies, it is readily apparent to those skilled in the art that the method may be used with other types of antibodies or with antibodies targetted against different

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epitopes or antigens. Other types of antibodies include but are not limited to fragments of antibodies and full-length antibodies.

The molecular biology and immunological techniques of the present invention can be performed by standard techniques well-known in the art. See, for example, in Maniatis, T., Fritsch, E.F., Sambrook, J., Molecular Cloning: A Laboratory Manual (Cold Spring Harbor Laboratory, Cold Spring Harbor, New York, 1982).

Because the genetic code is degenerate, more than one codon may be used to encode a particular amino acid, and therefore, the amino acid sequence can be encoded by any of a set of similar DNA oligonucleotides.

The cloned DNA molecules obtained may be expressed by cloning the gene encoding the altered antibody into an expression vector containing a suitable promoter and other appropriate transcription regulatory elements, and transferred into prokaryotic or eukaryotic host cells to produce recombinant modified antibodies. Techniques for such manipulations are well-known in the art.

In order to simplify the following Examples and the Detailed Description, certain terms will be defined.

Expression vectors are defined herein as DNA sequences that are required for the transcription of cloned copies of genes and the translation of their mRNAs in an appropriate host. Such vectors can be used to express eukaryotic genes in a variety of hosts such as bacteria, bluegreen algae, plant cells, insect cells and animal cells. Expression vectors include, but are not limited to, cloning vectors,

Expression vectors include, but are not limited to, cloning vectors, modified cloning vectors, specifically designed plasmids or viruses. Specifically designed vectors allow the shuttling of DNA between hosts, such as bacteria-yeast or bacteria-animal cells.

DNA encoding antibodies may also be cloned into an expression vector for expression in a host cell. Host cells may be prokaryotic or eukaryotic, including but not limited to bacteria, yeast, mammalian and insect cells and cell lines.

The expression vector may be introduced into host cells via any one of a number of techniques including but not limited to

transformation, transfection, protoplast fusion, and electroporation.

Expression of cloned DNA may also be performed using in vitro produced synthetic mRNA. Synthetic mRNA can be efficiently translated in various cell-free systems, including but not limited to wheat germ extracts and reticulocyte extracts, as well as efficiently translated in cell based systems, including but not limited to microinjection into frog oocytes, with micro-injection into frog oocytes being preferred.

It is also well-known that there is a substantial amount of redundancy in the various codons which code for specific amino acids. Therefore, this invention is also directed to those DNA sequences which contain alternative codons which code for the eventual translation of the identical amino acid. For purposes of this specification, a sequence bearing one or more replaced codons will be defined as a degenerate variant.

The following examples are provided to further define the invention without, however, limiting the invention to the particulars of these examples.

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EXAMPLE 1

Construction of mutations

Plasmid pP5Q was the starting vector for all mutagenic studies. Plasmid pP5Q is a derivative of p5H7 (Cambridge Antibodies). Plasmid pP5Q contains the VH and VL regions originally derived from MAb 447 (Gorney et al.) cloned as a single chain fragment variable (scFv).

Table 1 lists some of the oligonucleotide primers used for site-directed mutagenesis of complementary determining region 3 (CDR3) of MAb447. Primers were synthesized on either a model 381A DNA Synthesizer (Applied Biosystems, Foster City, CA) or a Cyclone™ Plus DNA Synthesizer (MilliGen/Biosearch, Marlborough, MA). Mutagenesis was performed with the Transformer™ Mutagenesis Kit (CLONTECH, Palo Alto, CA)

according to the manufacturer's instructions. All mutations were verified by DNA sequencing using the Sequenase® V2.0 DNA Sequencing Kit (United States Biochemical, Cleveland, OH).

Table 1

Primers:

Randomization of position 107: CTC GGA GAC TCC C/GNN AAT CAT AAA

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Randomization of position 111: GTA GTA GTC C/GNN GGA GAC TCC CCG

Randomization of position 112:
GTC GTT GTA GTA GTA GTA C/GNN CTC GGA GAC

EXAMPLE 2

Preparation of extracts and BIAcore analysis of scFv Extracts: 20 Mutagenized plasmids were introduced by electroporation into bacterial strain Escherichia coli TG1 for expression. Single colonies were inoculated into 10 ml of 2X-YT (which contains per liter of water 16 g tryptone, 10 g yeast extract and 5 g sodium chloride) supplemented with 2% glucose. Cells were 25 grown overnight at 30°C with vigorous shaking, collected by centrifugation in a Beckman GPR centrifuge at 2500 rpm, and resuspended in 10 ml of fresh 2X-YT supplemented with 1 mM isopropyl-beta-D-thiogalactopyranoside (IPTG) to induce expression. Cells were incubated at 30°C for an additional 5-6 hours with vigorous shaking, collected by centrifugation, resuspended in 1 ml of phosphate buffered saline: ethylenediametetraacetic acid (PBS:EDTA: 10 mM sodium phosphate pH7.0, 150 mM sodium chloride 1 mM EDTA), and incubated on ice for 30 minutes to

release periplasmic proteins. Extracts were clarified by centrifugation and stored at 4°C until use.

EXAMPLE 3

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Off-rate determinations of the scFv antibodies were determined using the BIAcore system (Pharmacia Biosenser). HIV gp120 V3 loop peptides, Al-1 variant (Ala-1 peptide) were covalently immobilized on a carboxylated dextran/gold matrix via the primary amino group. The carboxyl-dextran matrix was first activated with N-ethyl-N'-(3-diethylaminopropyl)carbondiimide (EDC) and reacted with N-hydroxysuccinimide (NHS). HIV gp120 V3 loop peptides such as Ala-1 peptide were covalently immobilized via the free thiol of a cysteine placed at the N-terminus. These peptides were reacted with the EDC-NHS activated matrix which had been reacted with 2-(2-pyridinyldithio)ethaneamine. Remaining unreacted NHS-ester groups were displaced by addition of ethanolamine. EDTA extracts were added in a flow passing over the immobilized antigen. The refractive index changes, in the form of the surface plasmon resonance caused by the binding and subsequent dissociation of the scFv, were monitored continuously. Off-rates were calculated from the automatically collected data using the Pharmacis Kinetics Evaluation software.

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EXAMPLE 4

Alanine scanning of CDR3 identifies residues which modulate scFvantigen_binding

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Alanine scanning mutagenesis was used to identify residues within the VH CDR3 region of scFv clone P5Q critical for binding. It was hypothesized that effects on binding by alanine substitution would lead to four broad classes of effect: class i) slower off-rate; class ii) faster off-rate; class iii) loss of binding; and class iv) minor or no change in off-rate. Class i) and ii) were

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operationally defined as critical. Class iii) was defined as obligatory. Class iv) was defined as noncritical.

The 27 positions that comprise VH CDR3 of scFv clone P5Q were individually changed to alanine by site-directed mutagenesis. Periplasmic extracts were prepared from the alanine replacement clones and assayed for off-rate determinations against the AL-1 gp120 V3 loop peptide (Fig. 1). Alanine substitutions at positions 107 and 111 resulted in 1.7 and 2.7 fold improvements in off-rate, respectively. These positions (class i) were judged critical and subsequently randomized to identify optimal residues. Alanine substitutions at positions 102, 112, 113, 114, and 118 led to faster off-rates (class ii); two of these positions were selected for further evaluation. Alanine substitution at positions 98, 101, 115, 116, 117, and 121 resulted in no binding (class iii). Alanine substitution at the remaining fourteen positions had only a minor effect on the off-rate (class iv). The class iii and iv positions were not evaluated further.

EXAMPLE 5

Randomization at critical positions to identify optimal amino acid solutions

The two critical class i) positions (107 and 111) were individually randomized to all amino acids, and off-rates against the AL-1 peptide determined. In addition, two class ii) positions (112 and 118) were also selected for randomization studies.

The results for position 107 are shown in Fig. 2. The slowest off-rate was observed with the negatively-charged glutamic acid, which decreased dissociation 2.5-fold. Substitution of other polar and charged amino acids had no significant effect on dissociation. With the exception of alanine, substitution with hydrophobic amino acid resulted in complete loss of binding. These results are consistent with the preponderance of surface ligand-contact residues being hydrophilic.

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Randomization of position 111 (Fig. 3) showed that the aromatic residues tyrosine and tryptophan produced the slowest off-rates (dissociation rates decreased 4.2 and 4.7-fold, respectively). However, substitution with any hydrophobic amino acids increased affinity relative to wild-type clone P5Q.

Class ii) positions 112 and 118 (faster off-rate upon alanine substitution) were also selected for amino acid randomization. For both position 112 (Fig. 4) and 118, the residues present in the original scFv P5Q, aspartic acid and asparagine, were the best solutions.

EXAMPLE 6

Improvements at positions 107 and 111 are additive

A double mutant that combined the optimized residues at positions 107 (E) and 111 (W) was constructed to determine whether or not the individual improvements are additive. Figure 5 shows that the double mutant has an off-rate 9-fold slower than wild-type clone P5Q. The off-rate value approximates the product of the fold improvements observed with the individual optimized residues (2.5 for 107E and 4.7 for 111W). One interpretation of this result is that for these two positions, the contributions to scFv-antigen affinity are independent and additive.

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EXAMPLE 7

Method of making modified antibodies

An antibody is mutagenized by alanine scanning mutagenesis to produce a modified antibody. The binding of the modified antibody to its antigen is determined. Binding determinations may be made by conventional methods and include off-rate measurements. Modified antibodies having desired characteristics are selected and maintained.

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EXAMPLE 8

Method of using modified antibodies

The modified antibodies or pharmaceutical compositions thereof are used for the prophylactic or therapeutic treatment of diseases caused by their antigen. Methods of treatment include, but are not limited to, intravenous or intraperitoneal injection of the modified antibody.

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EXAMPLE 9

Diagnostic kit employing modified antibodies

The modified antibodies of Example 7 are used as reagents in diagnostic kits. The modified antibody reagents may be further modified through techniques which are well-known in the art, such as radiolabeling or enzyme-labeling. The diagnostic kit may be used to detect or characterize the antigens.

EXAMPLE 10

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DNA encoding modified antibodies

The DNA encoding the modified antibody of Example 7 is used as a reagent for the production of modified antibodies. The DNA may be incorporated into an expression vector. The expression vector may be used to transform a host cell. Cultivation of the host cell under conditions suitable for the expression results in the production of modified antibody.

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EXAMPLE 11

DNA encoding modified antibodies

The DNA encoding the modified antibody of Example 7 is used to detect DNA encoding the antigen in test samples. Methods of detection include, but are not limited to, hybridization under selective conditions. Test samples include, but are not limited to, samples of blood, cells, and tissues.

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EXAMPLE 12

Preparation of modified light chain immunoglobulins

The light chain of an immunoglobulin is mutagenized by alanine scanning mutagenesis to produce a modified immunoglobulin having modified binding characteristics. The modified immunoglobulin is used as a reagent for diagnostic kits or as a therapeutic agent.

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SEQUENCE LISTING

(1) GENERAL INFORMATION:

- (i) APPLICANT: LEWIS, CRAIG M.
 LUDMERER, STEVEN W.
 HOLLIS, GREGORY F.
- (ii) TITLE OF INVENTION: IN VITRO ANTIBODY MATURATION
- (iii) NUMBER OF SEQUENCES: 2
- (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: CHRISTINE E. CARTY
 - (B) STREET: P.O. BOX 2000, 126 E. LINCOLN AVENUE
 - (C) CITY: RAHWAY
 - (D) STATE: NJ
 - (E) COUNTRY: USA
 - (F) ZIP: 07065
- (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25
- (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER: US 08/206,079
 - (B) FILING DATE: 04-MAR-1994
 - (C) CLASSIFICATION:
- (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: CARTY, CHRISTINE E.
 - (B) REGISTRATION NUMBER: 36,090
 - (C) REFERENCE/DOCKET NUMBER: 19190P
 - (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: (908) 594-6734
 - (B) TELEFAX: (908) 594-4720
- (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 816 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (genomic)

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(X1)	SEQUENCE	DESCRIPTION:	SEQ	ΤD	MO:I:

GC	CATGGCCG	AGGTGCAGCT	GGTGGAGTCT	GGGGGAGGCT	TGGTAAAGCC	TGGGGGGTCC	60
CT	CAGACTCA	CCTGTGTAGC	CTCTGGCTTC	ACGTTCAGTG	ATGTCTGGCT	GAACTGGGTC	120
CG	CCAGGCCC	CAGGGAAGGG	CCTGCACTGG	CTCCCCCCTA	TTAAAAGCGC	CACTGATGGT	180
GG	GACAACAG	ACTACGCTGC	ATCCGTGCAA	GGCAGATTCA	CCATCTCAAG	AGATGACTCA	240
AA	AAACACGC	TATATCTGCA	AATGAATAGC	CTGAAAACCG	AGGACACAGC	CGTTTATTCC	300
TG	CAACACAG	ATGGTTTTAT	TATGATTCGG	GGAGTCTCCG	AGGACTACTA	CTACTACTAC	360
AA	CGACGTTT	GGGGCAAAGG	GACCACGGTC	ACCGTCTCCT	CAGGTGCAGG	CGGTTCAGGC	420
GGJ	AGGTGGCT	CTGGCGGTGG	CGGATCGCAG	TCTGTGTTGA	CGCAGCCGCC	CTCAGTGTCT	480
GC	GCCCCAG	GACAGAAGGT	CACCATCTCC	TGCTCTGGAA	GCAGCTCCAA	CATTGGGAAT	540
AA:	PTATGTAT	TGTGGTACCA	GCAGTTCCCA	GGAACAGCCC	CCAAACTCCT	CATTTATGGC	600
AA'	TAATAA GC	GACCCTCAGG	GATTCCTGAC	CGATTCTCTG	GCTCCAAGTC	TGGCACGTCA	660
GC	CACCCTGG	GCATCACCGG	ACTCCAGACT	GGGGACGAGG	CCGATTATTT	CTGCGCAACA	720
TG	GATAGCG	GCCTGAGTGC	TGATTGGGTG	TTCGGCGGAG	GGACCAAGCT	GACCGTCCTA	780
GG:	rcccccc	CAGAACAAAA	ACTCATCTCA	GAAGAG			816

(2) INFORMATION FOR SEQ ID NO:2:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 272 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

Ala Met Ala Glu Val Glx Leu Val Glu Ser Gly Gly Gly Leu Val Lys 1 5 10 15

Pro Gly Gly Ser Leu Arg Leu Thr Cys Val Ala Ser Gly Phe Thr Phe 20 30

Ser Asp Val Trp Leu Asn Trp Val Arg Gln Ala Pro Gly Lys Gly Leu 35 40

Glu Trp Val Gly Arg Ile Lys Ser Ala Thr Asp Gly Gly Thr Thr Asp 50 60

Tyr 65	Ala	Ala	Ser	Val	Gln 70	Gly	Arg	Phe	Thr	Ile 75	Ser	Arg	Asp	Asp	Ser 80
Lys	Asn	Thr	Leu	Tyr 85	Leu	Glx	Met	Asn	Ser 90	Leu	Lys	Thr	G1u	Asp 95	Thr
Ala	Val	Tyr	Ser 100	Cys	Asn	Thr	Хар	Gly 105	Phe	Ile	Met	Ile	Arg 110	Gly	Val
Ser	Glu	Asp 115	Tyr	Tyr	Tyr	Tyr	Tyr 120	Asn	Asp	Val	Trp	Gly 125	Lys	Cly	Thr
Thr	Val 130	Thr	Ala	Ser	Ser	Gly 135	Ala	Gly	Gly	Ser	Gly 140	Gly	Gly	Gly	Ser
Gly 145	Gly	Gly	Ser	Gln	Ser 150	Val	Leu	Thr	Gln	Pro 155	Pro	Ser	Val	Ser	Ala 160
Ala	Pro	Gly	Gln	Lys 165	Val	Thr	Ile	Ser	Cys 170	Ser	Gly	Ser	Ser	Ser 175	Asn
Ile	Gly	Asn	Asn 180	Tyr	Val	Leu	Trp	Tyr 185	Gln	Gln	Phe	Pro	Gly 190	Thr	Ala
Pro	Lys	Leu 195	Leu	Ile	Tyr	Gly	Asn 200	Asn	Lys	Arg	Pro	Ser 205	Gly	Ile	Pro
Asp	Arg 210	Phe	Ser	Gly	Ser	Lys 215	Leu	Leu	Ile	Tyr	Gly 220	Ala	Thr	Leu	Gly
Ile 225	Thr	Gly	Leu	Gln	Thr 230	Gly	Asp	Gln	Ala	Asp 235	Tyr	Phe	Cys	Ala	Thr 240
Trp	Asp	Ser	Gly	Leu 245	Ser	Ala	Asp	Trp	Val 250	Phe	Gly	Gly	Gly	Thr 255	Lys
Leu	Thr	Val	Leu 260	Gly	Ala	Ala	Ala	Glu 265	Gln	Lys	Leu	Ile	Ser 270	Glu	Glu

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WHAT IS CLAIMED IS:

- 1. A DNA molecule encoding a modified antibody, the modified antibody being derived from a native antibody by alanine scanning mutagenesis and the modified antibody having binding characteristics different than binding characteristics of the native antibody.
- 2. The DNA molecule of Claim 1 wherein the native antibody is MAb447.
- 3. The DNA molecule of Claim 2, the DNA molecule being selected from the group consisting of P5Q, DNA encoding modified antibodies of Figures 1, 2, 3, 4, 5, combinations thereof, derivatives thereof and degenerate variants thereof.
 - 4. A method of modifying an antibody to make an modified antibody comprising replacing at least one amino acid of the antibody with alanine to produce a modified antibody.
 - 5. The method of Claim 4 wherein the modified antibody has improved binding characteristics.
- 6. Modified antibodies produced by the method of Claim 4 or homologues thereof.
 - 7. The method of Claim 4 wherein the antibody is MAb447.
- 8. The method of Claim 7 wherein the amino acid replaced with alanine is located in complementary determining region 1, complementary determining region 2 or complementary determining region 3 of MAb447.

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- 9. The modified antibodies of Claim 6 selected from the group consisting of P5Q, the antibodies of Figures 1, 2, 3, 4, 5, combinations thereof, derivatives thereof, and homologues thereof.
- 10. Diagnostic kits comprising the modified antibodies produced by the method of Claim 6.
- 11. Diagnostic kits comprising the DNA molecules of Claim 1.
- 12. A pharmaceutical composition comprising at least one modified antibody of Claim 6 or DNA encoding at least one modified antibody of Claim 6 or combinations thereof.

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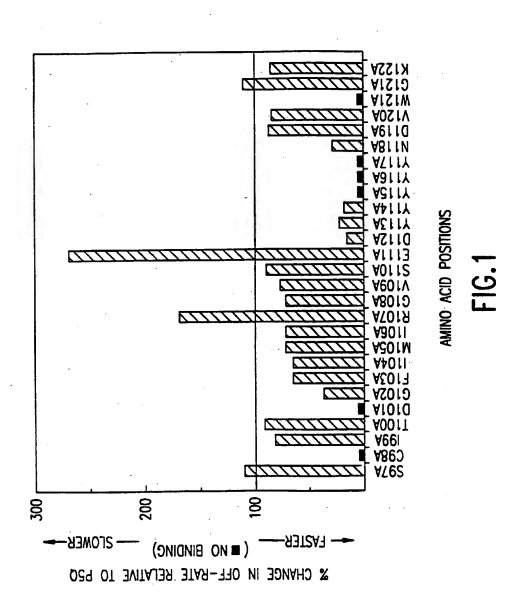
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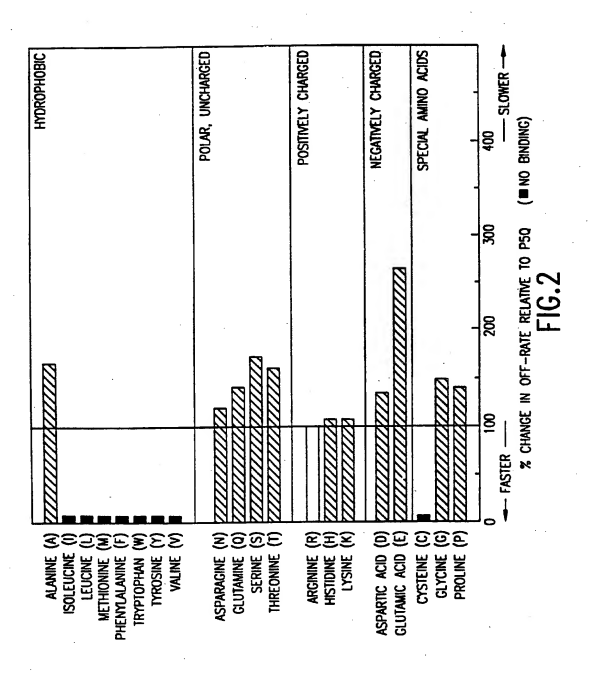
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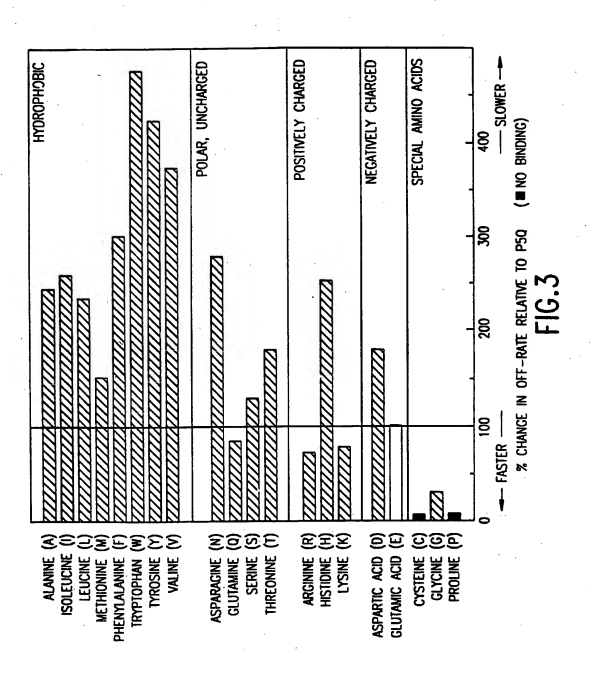
20

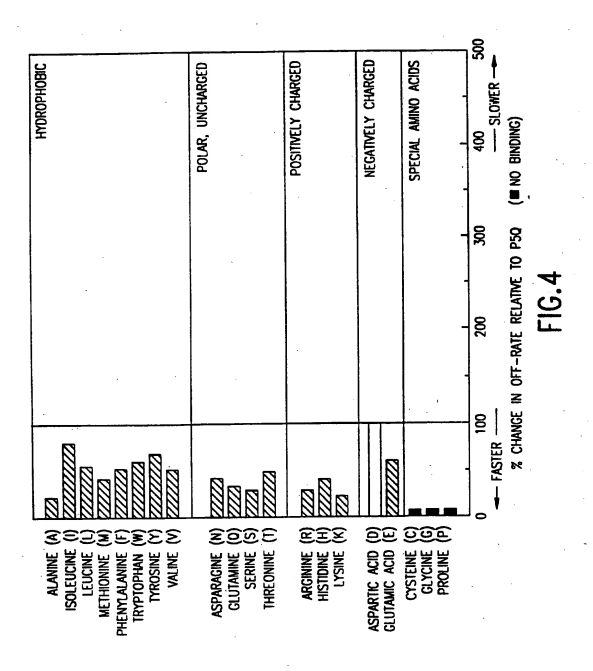
25

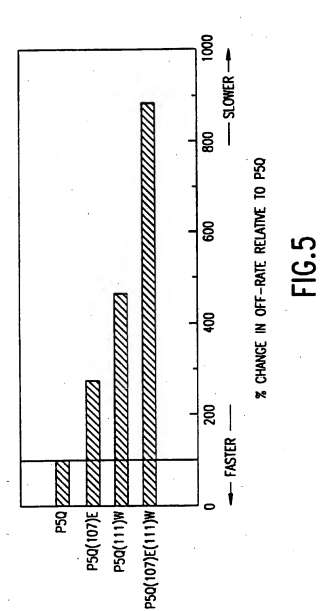
30











									6	/9					
9	*	TCC	Ser	120	#	GTC	Val	180	#	GGT	(4)	240	٠	TCA	Sr
. •		2000	Gly			TGG	Trp			GAT				GAC	Asp
		999	Gly			AAC	Asn '			ACT			:	CAT	Asp
20	*	CCI		110	#	CTG		170	*	GCC	t	230	•	AGA	Arg
			Lys	H		TGG		+		AGC		7		TCA	Ser
		GTA	Val			GTC	Val			AAA Ive			•	ATC	Ile
0		•	Leu	0		GAT		0	*	ATT) 4	0	*	ACC	Thr
4			Gly	100		AGT	Ser	160		CGT		220		TTC	Phe
			Gly (TTC				ည္ပင္မ					Arg
		999	Gly			ACG				GTC					Gly
30	•	TCT		90	*		Phe	150	#	766		210	•	CAA	Gln
		GAG	Gla			ည္သ	Gly			GAG				GTG	Val
		GTG	Val			TCT	Ser			CTG				TCC	Ser
20	#1	CTG	Leu	80	#	ညည		40	#	666		200	•		Ala
		CAG		-		GTA	Val	. .		AAG		7		CCT	Ala
							Cys				7		٠		
0	•	GAG	Glu	70		ACC	Thr	130		CCA GGG) 	0		GAC	Asp
_		ညည	Ala	7		CTC	Leu	13		GCC		190		ACA	Thr
		ATG	Ala Met Ala Glu Val			AGA CTC ACC	Arg			CGC CAG GCC CCA GGG				ACA	Thr Thr Asp Tyr
		CCC	Ala			CIC /	Leu		•	CGC	N			999	Gly .

FIG.60

				7/9	
300	TCC	360	TAC	420 * GGC G1y	480 TCT Ser
	TAT		TAC	TCA	GTC Val
	GTT Val		TAC	GGT	TCA
290	GCC	350	TAC	410 GGC Gly	470 CCC Pro
73	ACA	מו	TAC	GCA	CCG
	GAC		GAC	GGT	460 TTG ACG CAG
O *	GAG	0 *	GAG	o TCA Ser	ACG
280	Acc Thr	.340	TCC GAG Ser Glu	400 t TCC TCA Ser Ser	460 TTC AC
	AAA Lys		GTC Val	GTC	GTG
	CTG	•	GGA Gly	ACC	TCT
270	AGC	330	CGG	390 * GTC Val	450 * CAG
	AAT Asn		ATT 11e	ACG	TCG
	ATG Met		ATG Met	Acc Thr	GGA
260	CAA	320	ATT Ile	380 GGG G1y	440 440 GGC G1v
~	CTG	m ,	TTT Phe	AAA Lys	GGT V
	TAT		GGT Gly	GGC	GGC
0 +	CTA	310	AAC ACA GAT GG Asn Thr Asp Gl)	370 * T TGG GGG	430 * C TCT GGG
250	ACG	31	ACA Thr	37 GTT Val	4. GGC V
	AAA AAC ACG CTA TAT Lys Asn Thr Leu Tyr			370 * AAC GAC GTT TGG GGC ASn ASP Val Trp Gly	43 GGA GGT GGC
	AAA		TGC Cys	AAC	GGA

FIG. 6b

540	• E	Ę	Asn	600	*	ပ္ပပ္ပ	Gly	7	9	•	TCA	Ser	720	•	ACA	Thr
	ָ נ	3	Gly			TAT	Tyr				ACG	Thr			ຽ	Ala
		7 7 4	I:le				Ile				gg	Gly			TGC	Cys
530	ינ בייל בייל	ر ج	Asn	590	*	CIC	Leu	650	2 .	•	TCT	Ser	710	*	TTC	Phe
ഗ	Ç	ر د	Ser	w		CIC	Leu	ų)		₽	Lys	•		AT	χŁ
	(Ser			AAA	Pro Lys			•	TCC	Gly Ser 1			GAT	p Glu Ala Asp T
o (ָ ֡ ע	200	Ser	0	*	ပ္ပပ္ပ	Pro	c) +	•	ပ္ပပ္ပ	Gly	0		ပ္ပပ္ပ	Ala
520	ξ	5	Gly	580		ပ္ပ	Ala	640			Ţ	Ser	700		GAG	GJn
	ي ا	֚֚֚֚֚֚֚֚֚֡֝֝֝֝֝֟֝֝֟֝֓֓֓֓֓֓֓֓֓֓֓֡֓֓֓֓֓֡֓֡֓֡֓֡֓֡֡֡֡֡֓֓֡֡֡֡֡֡֡	Ser			ACA	Gly Thr				TTC	Phe	•		GAC	Gly Asp (
	נ נ	נ פ	Cys			GGA	Gly				CGA	Arg		٠	၁၁၁	Gly
510	. ر د		Ser	570	*	CCA	Pro	630)	S	GAC	Asp	069	*	ACT	Thr
			Ile			TTC	Phe				CCT	Pro				Gln
	ن د د		Thr			CAG	Gln				ATT	Ile		,	CTC	ren
200	. į	ָ כלי כלי	Val	260	•	CAG	Gln	620) (ပ္ပပ္ပ	Gly	089	•	GGA	Gly
u 1	()	2	Lys			TAC	Tyr	•	•		TCA	Ser			ACC	Thr
	ć	כ כ	Gln			TGG	Trp				ပ္ပ	Pro	٠.		ATC ACC	Ile
490	֓֞֝֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓	5	Gly	550	•	TTC	Leu	610) 1		CGA	Arg	670	*	ညည	Gly
4	ξ	נ נ	Pro	Ŋ		TAT GTA 1	Val	Ý	ò		AAG	Lys	io		CTC	Len
	ָ נ	ر ر	Ala	-		TAT	Tyr				AAT AAT AAG CGA	Asn			S	먑
	į	ر د د	Ala			AAT	Asn				AAT	Asn			ညည	Ala

FIG. 60

			•
780	CTA		
	GTC		
	ACC		
770	CTG Leu		
	AAG		
	ACC		
0 *	GGG		
760	GGA		din.
	GGC Gly		GAG
	TTC		GAA
750	GTG TTC	810	Ser
	TGG		AIC
	GAT		GAA CAA AAA CTC ATC TCA GAA GAG Glu Gln Lys Leu Ile Ser Glu Glu
740	CTG AGT GCT GAT Leu Ser Ala Asp	800	AAA
	AGT		Gla
	CTG		
730	GGC	190	GCA Ala
7	AGC	7	GCC
	GAT ASP		GCG
	TGG (Trp		GGT

FIG. 6d